## **PCT**

# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



C07C 29/136, 31/20 A1 21) International Application Number: PCT/EP99/02 22) International Filing Date: 2 April 1999 (02.04.	BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB,
22) International Filing Date: 2 April 1999 (02.04.	BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB,
9800276  9 April 1998 (09.04.98)  71) Applicant (for all designated States except US): PANTOCH S.A. [BE/BE]; Parc Industriel, Zone A, B-7181 Feluy (E)  72) Inventors; and  75) Inventors/Applicants (for US only): BERTOLA, Aldo [IT/Via Luigi Illica, 5, I-20121 Milano (IT). RANGHII Giovanni [IT/IT]; Via 7a Strada, 27, I-20090 San lice-Segrate (IT).  74) Agent: SARPI, Maurizio; Studio Ferrario, Via Collina, I-00187 Roma (IT).	KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU TM), European patent (AT, BE, CH, CY, DE, DK, ES FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI pt (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, SN, TD, TG).  Published  With international search report.  Before the expiration of the time limit for amending claims and to be republished in the event of the receip

Process for the production of 1,4–Butanediol by vapour phase catalytic hydrogenation of gamma–butyrolactone, maleic and/or succinic anhydride methyl esters or mixtures of gamma–butyrolactone and maleic and/or succinic anhydride esters, characterised by the fact that, in the operating conditions described in the process, by feeding hydrogen gas containing carbon monoxide to the reaction, a concomitant production of synthesis methanol is obtained, and conversion values relating to methanol production are improved.

### FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA.	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
ВЈ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
СН	Switzerland	KG	Kyrgyzstan	NO	Norway	ZW	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
cz	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	$\mathbf{s}\mathbf{G}$	Singapore		

#### Process for the production of Butanediol.

10

15

20

#### Description.

The present invention relates to a process for the production of 1,4-butanediol by vapour phase flow catalytic hydrogenation of gamma-butyrolactone, succinic or maleic anhydride methyl esters, or mixtures thereof, wherein the hydrogen flow is enriched with CO, so as to accomplish an extra production of synthesis methanol, with a consequent increase in the yield of the final product.

It is known from the prior art that there exist many processes for the preparation of butanediol using dicarboxylic acid esters with four carbon atoms as starting materials.

In USSR Patent  $N^{\circ}400,567$  a copper chromite catalysed liquid phase conversion of esters into BDO at temperatures ranging between 280 and 300°C and 300 bar pressure is given.

In US Patent  $N^4$ ,613,707 a conversion of diethylsuccinate into BDO at 200°C T and 135 bar pressure with a mainly copper and aluminium borate catalyst is given.

In US Patent  $N^{\circ}2,079,414$  a vapour phase hydrogenation of esters is given, on a copper chromite type catalyst and temperatures ranging between 300 and  $400^{\circ}\text{C}$ .

In US Patent n°2,040,944 a broad description of both liquid and vapour phase ester hydrogenations is given, using copper chromite catalysts. In one example, a liquid phase butyl succinate hydrogenation at 207 bars is described.

In US Patents n°'s 4,584,419 and 4,751,334, assigned to Davy Mc Kee Ltd. (U.K.) processes for the large scale production of BDO are described and they involve four carbon dicarboxylic acid ester hydrogenations.

The operating conditions of the hydrogenations claimed in the patents assigned to Davy McKee Ltd. are pressures ranging between 25 and 75 bars, and temperatures between 150° and 240°C, in the presence of a stabilised cuprous chromite type catalyst in its reduced form.

Moreover in the latter patent it is stated that the process which is claimed in it yields a 79.3% conversion into the desired product, after the first stage.

There is no indication of the overall conversion, but it is not at all unfair to be sure that it is somewhat lower than the above value.

Likewise, a 78.3% overall conversion is obtained in the process claimed in the former patent.

Clearly, as the above values hold on an industrial scale, they leave much room for substantial improvements.

Aim of the present invention is therefore to propose a process which allows much higher conversion rates, therefore increases the profitability of its application to the plant scale.

Another aim of the present invention is that of proposing a methodology for the contemporaneous production of BDO and methanol (MOH) operating in specific conditions and on suitable catalysts.

The above aims are accomplished by a process for the production of 1,4-butanediol by vapour phase selective hydrogenation of gamma-butyrolactone, maleic or succinic anhydride methyl esters or mixtures of gamma-butyrolactone with the esters thereof, characterised by the fact that synthesis methanol is coproduced by feeding hydrogen gas enriched with carbon monoxide to the reaction.

According to the process of the present invention, methyl dicarboxylic acid esters with four carbon atoms, e.g. dimethylmaleate (DMM) or dimethylsuccinate (DMS), gamma butyrolactone (GBL), or mixtures of GBL with the esters thereof, are utilised as starting materials for the production of BDO.

Methanol synthesis is accomplished by feeding carbon monoxide together with hydrogen gas. The reaction by which BDO forms thanks to the hydrogenation of an ester like DMS for example, is given by the following equation:

- (A)  $CH_3-O-CO-CH_2-CH_2-CO-O-CH_3+4H_2\rightarrow 2CH_3OH+OH(CH_2)_4OH$  The MOH synthesis reaction is given by the following equation:
  - (B)  $CO+2H_2\rightarrow CH_3OH$

MOH that forms in reaction (A) is recovered to be fed to an esterification unit .

However from a practical viewpoint, part of the MOH is lost either literally or in the form of by-products. MOH real losses, about 0.1 to 0.3 mols for each mol of BDO produced, may even be compensated by MOH formation according to reaction (B), this making the production unit completely self-sufficient in terms of MOH.

The most important aspect is the neat improvement given by the process object of the present invention in so far as the yield of reaction (A) is concerned, more than its self-sufficiency in terms of MOH consumption, the latter being nevertheless a not unimportant factor.

Surprisingly, the MOH synthesised which adds up to the MOH produced in reaction (A) is not of any hindrance to the BDO synthesis reaction as it would be expected to be. Instead it results to have a remarkably positive effect on the conversion of the ester and/or GBL of the shot.

The surprising observation stated above may be explained by dint of the protective action exerted onto the active sites of the catalyst by the methanol synthesis.

It may be significant that the catalyst employed in the hydrogenation reaction is of either the exact same or similar type as that employed in methanol synthesis, e.g. of the copper-zinc oxide, or cuprous chromite type, this ensuring a compatibility and ease of coexistence of the two reactors in the same vessel.

In the process which is object of the present invention, the typical operating conditions for this reaction are as follows:

Reaction type:

adiabatic

Molar ratio between H2 and ester:

from 200 to 800,

preferably from

200 to 500

Molar ratio between CO and

shot ester: from 0.1 to 1.0

preferably from

0.2 to 0.4

Operating pressure: from 75 to 120

bars, preferably

from 80 to 100

bars

Operating temperature: from 170 to 230°C

preferably from

190° to 210 °C

Type of catalyst: Copper/Zinc oxide

or cuprous

chromite

Liquid Hourly Space Velocity: from 0.1 to 1.0

hr<sup>-1</sup>, and

averagely from

0.3 to 0.5  $hr^{-1}$ 

These and other features will be more readily apparent from the following description of a preferred not limiting embodiment of the invention with reference to the accompanying drawing in which a scheme (Fig. 1) of the production process is shown.

The following operating conditions refer to a starting material made up of a mixture of GBL and DMS in 70:30 proportions by weight.

The feed mixture (Line 1) is fed to vapouriser 3, together with a liquid recycle mixture (Line 2), that contains GBL from fractionation unit 22. In vapouriser 3 both the feed (Line 1) and the recycle (Line 2) come to contact with a hot hydrogen stream (Line 4), and they vapourise. In the gas stream coming out of the vapouriser (Line 5) the molar ratio between hydrogen and ester is 300:1, the molar ratio between CO and ester is 0.2, temperature is 190°C and pressure is 80 ATE.

Such stream feeds Stage I of reaction 6 where there is a copper-zinc oxide type catalyst with a surface area never any smaller than 40  $\rm m^2g^{-1}$ .

At the inlet of Stage II in reaction 7, temperature is taken down to approx. 190°C by injection of a cold hydrogen stream (Line 7). Overall Liquid Hourly Space Velocity in the two reaction stages is 0.35  $hr^{-1}$ .

At the inlet of Stage II in reaction 8, DMS conversion results to be higher than 98%, while GBL conversion in the feed is higher than 91%.

The overall average conversion value works out to be as high as 93%.

By-products obtained in the process consist of tetrahydrofuran (THF)-about 6% on a molar base with respect to the BDO produced.

Selectivity of the reaction is approx. 99% with respect to the overall amount of BDO and THF produced.

The high molar ratio between hydrogen and carbon monoxide favours the occurrence of methanol synthesis with stepwise conversion values higher than 90%.

The effluent from the reactor (Line 9) cools down in exchanger 10, letting heat to the recycle hydrogen rich stream, and in exchanger 11. Eventually it feeds (Line 20) separator 12 where the condensed organic phase separates from the hydrogen rich gaseous phase.

The gaseous phase coming out of separator 12 (Line 13) is compressed by compressor 14 to be recycled into the reaction system.

A small fraction of the recycle gas is purged (Line 15) to minimise the accumulation of inert materials.

The compressed gas (Line 16) and the feed hydrogen and carbon monoxide (Line 17), partly (Line 7) blend with the effluents from the first stage of reaction 6, and partly (Line 18) pre-heat in exchanger 10 and in terminal heater 19, to subsequently feed (Line 4) vapouriser 3.

The liquid phase coming out of separator 12 feeds (Line 21) product fractionation unit 22, where THF (Line 23), MOH (Line 24), water and light organic by-

products (Line 25), an unconverted GBL rich organic fraction (Line 2) bound to be recycled for hydrogenation, heavy organic by-products (Line 26) and BDO (Line 27) separate.

The innovative aspect of the process object of the present invention is better emphasised by a comparison of its performances of which above with those obtained carrying out the hydrogenation with the same shot and operating conditions, nevertheless without streaming any carbon monoxide gas.

It was found that without carbon monoxide, hence without any MOH synthesis, keeping the same high selectivity values, the overall conversion of GBL to DMS decreases from the 93% mark, obtained when feeding carbon monoxide, to approx. 84 %.

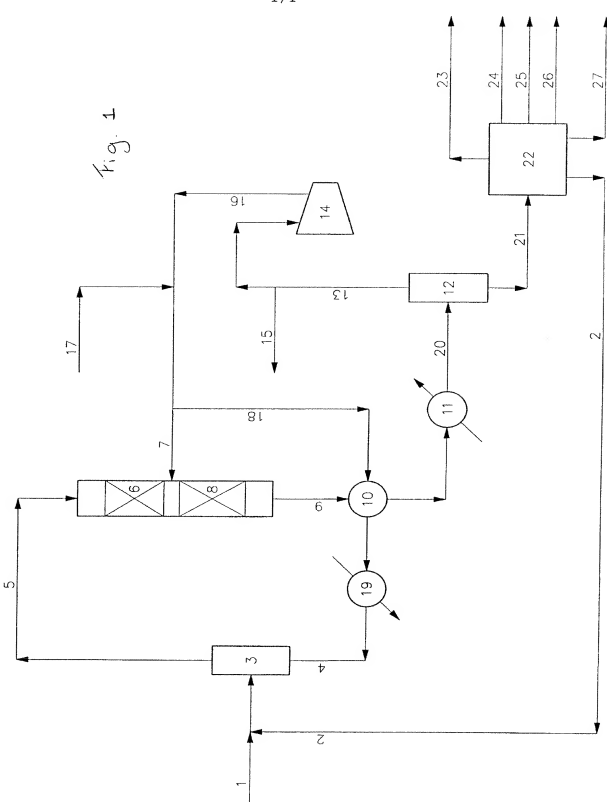
The lower conversion rates entice higher investment and production costs, both in the hydrogenation section, and in the subsequent distillation sections where the unconverted fractions bound to be recycled for hydrogenation are separated.

#### CLAIMS

- 1. A process for the production of 1,4-butanediol by vapour phase selective hydrogenation of gamma-butyrolactone, maleic or succinic anhydride methyl esters or mixtures of gamma-butyrolactone with the esters thereof, characterised by the fact that synthesis methanol is coproduced by feeding hydrogen gas enriched with carbon monoxide to the reaction.
- 2. A process according to claim 1, characterised in that the molar ratio between carbon monoxide and shot ester ranges between 0.1 and 1.0
- 3. A process according to claims 1 and 2, characterised in that the hydrogen to shot ester molar ratio ranges between 200 and 800.
- 4. A process according to claims 1,2 and 3, characterised in that its operating pressure ranges between 75 and 100 bars, and its operating temperature ranges between 170° and 230°C.
- 5. A process according to claims 1,2,3 and 4, characterised in that the hydrogenation catalyst is of the copper-zinc oxide or stabilised cuprous chromite type, with surface area never any smaller than  $40~\text{m}^2\text{g}^{-1}$ .
- 6. A process according to claim 5, characterised in that the vapour phase mixture containing shot hydrogen and ester contacts the catalyst

with a liquid hourly space velocity ranging between 0.1 and 1.0  $hr^{-1}$ .





# INTERNATIONAL SEARCH REPORT

rnational Application No PCT/EP 99/02297

		PC	1/EP 99/0229/
A. CLASSI IPC 6	FICATION OF SUBJECT MATTER C07C29/136 C07C31/20		
According to	o International Patent Classification (IPC) or to both national classif	ication and IPC	
	SEARCHED		
Minimum do	ocumentation searched (classification system followed by classification sy	ation symbols)	
1100	0070		
Documental	tion searched other than minimum documentation to the extent that	such documente are included in	n the folds searched
	and the second state with the second state of	. oddir doddinerka are molddd i	in the helds seatched
Electronic di	ata base consulted during the international search (name of data b	ease and where practical searc	h tarme ((sad)
	,	and more presently source	Tromis useuj
C. DOCUME	ENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the r	elevant passages	Relevant to claim No.
Υ	US 4 584 419 A (SHARIF MOHAMMAD	ET AL)	1-6
	22 April 1986 (1986-04-22) cited in the application		
	column 4, line 42 - line 44; cl	aim 1;	
	example 1; table IV		
Υ	EP 0 722 923 A (STANDARD OIL CO	OHIO)	1-6
	24 July 1996 (1996-07-24)	,	
	page 3, line 19 - line 21; clai	ms 1-3	
			1
1			
Furth	er documents are listed in the continuation of box C.	X Patent family membe	rs are listed in annex.
° Special cat	egories of cited documents :	"T" later document published a	fter the international filing date
"A" documer conside	nt defining the general state of the art which is not pred to be of particular relevance	or priority date and not in cited to understand the pr	conflict with the application but inciple or theory underlying the
	ocument but published on or after the international	invention "X" document of particular rele	
which is	nt which may throw doubts on priority claim(s) or s cited to establish the publication date of another	involve an inventive step v	el or cannot be considered to when the document is taken alone
citation "O" docume	or other special reason (as specified)  nt referring to an oral disclosure, use, exhibition or	"Y" document of particular rele- cannot be considered to in document is combined wit	vance; the claimed invention  Notice an inventive step when the  h one or more other such docu-
other m	eans  t published prior to the international filing date but	ments, such combination l in the art.	being obvious to a person skilled
later tha	an the priority date claimed	"&" document member of the sa	
Date of the a	ctual completion of the international search	Date of mailing of the inter	national search report
11	August 1999	18/08/1999	
Name and ma	ailing address of the ISA	Authorized officer	
	European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk		
	Tel. (+31-70) 340-2040. Tx. 31 651 epo ni, Fax: (+31-70) 340-3016	Arias-Sanz,	J

### INTERNATIONAL SEARCH REPORT

Information on patent family members

rnational Application No
PCT/EP 99/02297

Patent document cited in search report		Publication date	•	Patent family member(s)	Publication date
US 4584419	Α	22-04-1986	CA EP JP	1246616 A 0143634 A 61022035 A	13-12-1988 05-06-1985 30-01-1986
EP 0722923	A	24-07-1996	US AT CN DE DE ES JP	5473086 A 178878 T 1129206 A 69509060 D 69509060 T 2130538 T 8193040 A	05-12-1995 15-04-1999 21-08-1996 20-05-1999 05-08-1999 01-07-1999 30-07-1996